What is claimed is:

 A method for comparing a first microarray spot with a second microarray spot comprising:

providing a first plurality of intensity values ( $S_i^A$ ) for said first microarray spot and a second plurality of intensity values ( $S_k^B$ ) for said second microarray spot;

calculating a p value using Wilcoxon's rank sum test, wherein said p value is for a null hypothesis that  $\theta=0$  and an alternative hypothesis that said  $\theta>0$ , wherein said  $\theta$  is a test statistic for intensity difference between said first plurality and said second plurality; and

indicating said first microarray spot is different from said second microarray spot if said p value is greater than a significance level.

- 2. The method of Claim 1 wherein said testing statistic is median  $(S_k^A)$ -median  $(S_k^B)$ .
- 3. The method of Claim 2 wherein said significance level is 0.05.
- 4. The method of Claim 1 wherein said first microarray spot and second microarray spot are nucleic acid spots.
- 5. The method of Claim 4 wherein said nucleic acid spots are among at least 100 nucleic acid spots on a substrate.

- The method of Claim 5 wherein said nucleic acid spots are among at least 1000 spots on said substrate.
- 7. The method of Claim 6 wherein said nucleic acid spots are cDNA spots.
- 8. The method of Claim 7 wherein said nucleic acid spots are oligonucleotide spots.
- 9. The method of Claim 1 further comprising step of combining first plurality and second plurality of intensity values if said *p*-value is greater than a significance level.
- 10. A computer software product for comparing a first microarray spot with a second microarray spot comprising:

computer program code for inputing a first plurality of intensity values  $(S_i^A)$  for said first microarray spot and a second plurality of intensity values  $(S_k^B)$  for said second microarray spot;

computer program code for calculating a p value using Wilcoxon's rank sum test, wherein said p value is for a null hypothesis that  $\theta=0$  and an alternative hypothesis that said  $\theta>0$ , wherein said  $\theta$  is a test statistic for intensity difference between said first plurality and said second plurality; and

computer program code for indicating said first microarray spot is different from said second microarray spot if said p value is greater than a significance level; and

a computer readable media for storing said computer program codes.

- 11. The computer program product of Claim 10 wherein said testing statistic is  $median(S_k^A)$ -median( $S_k^B$ ).
- 12. The computer program of Claim 11 wherein said significance level is 0.05.
- 13. The computer software product of Claim 11 further comprising computer program code for accepting user's input or selection of said significance level.
- 14. The computer software product of Claim 11 wherein said first microarray spot and second microarray spot are nucleic acid spots.
- 15. The computer software product of Claim 14 wherein said nucleic acid spots are among at least 100 nucleic acid spots on a substrate.
- 16. The computer software product of Claim 15 wherein said nucleic acid spots are among at least 1000 spots on said substrate.

- 17. The computer software product of Claim 16 wherein said nucleic acid spots are cDNA spots.
- 18. The computer software product of Claim 16 wherein said nucleic acid spots are oligonucleotide spots.
- 19. The computer software product of Claim 10 further computer program code for combining first plurality and second plurality of intensity values if said *p*-value is greater than a significance level.
- 20. The computer software product of Claim 19 wherein said significance level is 0.5.
- 21. A system for comparing nucleic acid probes, comprising:
  - a processor; and

a memory being coupled to the processor, the memory storing a plurality machine instructions that cause the processor to perform a plurality of logical steps when implemented by the processor, said logical steps including:

inputing a first plurality of intensity values  $(S_i^A)$  for said first microarray spot and a second plurality of intensity values  $(S_k^B)$  for said second microarray spot;

calculating a p value using Wilcoxon's rank sum test, wherein said p value is for a null hypothesis that  $\theta$ =0 and an alternative hypothesis that said  $\theta$ >0,

wherein said  $\theta$  is a test statistic for intensity difference between said first plurality and said second plurality; and

indicating said first microarray spot is different from said second microarray spot if said p value is greater than a significance level.

- 22. The system of Claim 21 wherein said testing statistic is *median* ( $S_i^A$ )
  median( $S_k^B$ ).
- 23. The system of Claim 22 wherein said significance level is 0.05.
- 24. The system of Claim 22 wherein said steps further comprise accepting user's input or selection of said significance level.
- 25. The system of Claim 21 wherein said first microarray spot and second microarray spot are nucleic acid spots.
- 26. The system of Claim 25 wherein said nucleic acid spots are among at least 100 nucleic acid spots on a substrate.
- 27. The system of Claim 26 wherein said nucleic acid spots are among at least 1000 spots on said substrate.

- 28. The system of Claim 27 wherein said nucleic acid spots are cDNA spots.
- 29. The system of Claim 27 wherein said nucleic acid spots are oligonucleotide spots.
- 30. The system of Claim 21 wherein said steps further comprise combining first plurality and second plurality of intensity values if said *p*-value is greater than a significance level.
- 31. The system of Claim 30 wherein said significance level is 0.5.
- 32. A method for determining whether a transcript is present in a biological sample comprising:

providing a plurality of perfect match pixel intensity values  $(PM_{ij})$  and mismatch pixel intensity values  $(MM_{ik})$  for the transcript, wherein said  $PM_{ij}$  is the pixel intensity value for perfect match probe i and pixel j and  $MM_{ik}$  is the pixel intensity value for mismatch probe i and pixel k;

calculating a p-value using one-sided Wilcoxon's rank sum test, wherein the p-value is for a null hypothesis that  $(median(PM_{ij})-median(MM_{ik}))=a$  threshold value and an alternative hypothesis that said  $(median(PM_{ij})-median(MM_{ik}))>$  said threshold value; and

indicating whether said transcript is present based upon said p-value.

- 33. The method of Claim 32 wherein said threshold value is zero.
- 34. The method of Claim 32 wherein said threshold value is calculated using:  $\tau = c \sqrt{median(PM_1)}$  wherein said c is a constant.
- 35. The method of Claim 32 wherein threshold value is calculated using:  $\tau = c_1 \sqrt{mean(PM_i)}$  wherein said c is a constant.
- 36. The method of Claim 32 wherein said step of indicating comprises indicating said transcript is present if said p is smaller than a first significance level  $(\alpha_1)$ .
- 37. The method of Claim 32 wherein said step of indicating further comprises indicating said transcript is absent if said p is greater than or equal to a second significance level  $(\alpha_2)$ .
- 38. The method of Claim 37 wherein said step of indicating further comprises indicating said transcript is marginally detected if  $\alpha_1 \le p < \alpha_2$ .
- 39. A computer software product for determining whether a transcript is present in a biological sample comprising:

computer program code for inputting a plurality of perfect match pixel intensity values  $(PM_{ij})$  and mismatch pixel intensity values  $(MM_{ik})$  for said transcript, wherein said  $PM_{ij}$  is the pixel intensity value for perfect match probe i and pixel j and  $MM_{ik}$  is the pixel intensity value for mismatch probe i and pixel k;

computer software code for calculating a p-value using one-sided Wilcoxon's rank sum test, wherein the p-value is for a null hypothesis that  $(median(PM_{ij})-median(MM_{ik}))=$ a threshold value and an alternative hypothesis that said  $(median(PM_{ij})-median(MM_{ik}))>$  said threshold value;

computer software code for indicating whether said transcript is present based upon said p-value; and

a computer readable media for storing said code.

- 40. The computer software product of Claim 32 wherein said threshold value is zero.
- 41. The computer software product of Claim 32 wherein said threshold value is calculated using:  $\tau = c \sqrt{median(PM_1)}$  wherein said c is a constant.
- 42. The computer software product of Claim 32 wherein threshold value is calculated using:  $\tau = c_1 \sqrt{mean(PM_1)}$  wherein said c is a constant.

- 43. The computer software product of Claim 32 wherein said computer program code for indicating comprises computer software code for indicating that said transcript is present if said p is smaller than a first significance level  $(\alpha_1)$ .
- 44. The computer software product of Claim 32 wherein said computer program code for indicating further comprises computer software code for indicating said transcript is absent if said p is greater than or equal to a second significance level  $(\alpha_2)$ .
- 45. The computer software product of Claim 37 wherein said computer program code for indicating further comprises computer software code for indicating that said transcript is marginally detected if  $\alpha_1 \le p < \alpha_2$ .
- A system for comparing nucleic acid probes, comprising:
  a processor; and
  a memory being coupled to the processor, the memory storing a plurality machine
  instructions that cause the processor to perform a plurality of logical steps when
  implemented by the processor, said logical steps including:

providing a plurality of perfect match pixel intensity values  $(PM_{ij})$  and mismatch pixel intensity values  $(MM_{ik})$  for the transcript, wherein said  $PM_{ij}$  is the pixel intensity value for perfect match probe i and pixel j and  $MM_{ik}$  is the pixel intensity value for mismatch probe i and pixel k;

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calculating a p-value using one-sided Wilcoxon's rank sum test, wherein the p-value is for a null hypothesis that  $(median(PM_{ij})-median(MM_{ik}))=a$  threshold value and an alternative hypothesis that said  $(median(PM_{ij})-median(MM_{ik}))>$  said threshold value; and

- 5 indicating whether said transcript is present based upon said *p*-value.
  - 47. The system of Claim 46 wherein said threshold value is zero.
  - 48. The system of Claim 47 wherein said threshold value is calculated using:  $\tau = c \sqrt{median(PM_i)}$  wherein said c is a constant.
    - 49. The system of Claim 47 wherein threshold value is calculated using:  $\tau = c_1 \sqrt{mean(PM_t)}$  wherein said c is a constant.
- 15 50. The system of Claim 46 wherein said step of indicating comprises indicating said transcript is present if said p is smaller than a first significance level  $(\alpha_1)$ .
  - 51. The system of Claim 50 wherein said step of indicating further comprises indicating said transcript is absent if said p is greater than or equal to a second significance level  $(\alpha_2)$ .

52. The system of Claim 51 wherein said first significance level  $(\alpha_1)$  is smaller than said  $(\alpha_2)$  and said step of indicating further comprises indicating said transcript is marginally detected if  $\alpha_1 \le p < \alpha_2$ .

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